

RN 128273-78-9 CAPLUS  
CN Thiourea, N-[[4-[(bromodifluoromethyl)thio]phenyl]methyl]-N'-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

[--00000106]

RN 128273-79-0 CAPLUS  
CN Thiourea, N-[[4-(heptafluoropropyl)phenyl]methyl]-N'-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

[--00000107]

RN 128273-80-3 CAPLUS  
CN Thiourea, N-[[4-[3,3,4,4,5,5,5-heptafluoro-2,2-bis(trifluoromethyl)pentyl]phenyl]methyl]-N'-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

[--00000108]

RN 128273-81-4 CAPLUS  
CN Thiourea, N-[[4-(tridecafluorohexyl)phenyl]methyl]-N'-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

[--00000109]

RN 128273-84-7 CAPLUS  
CN Thiourea, N-[[4-[[3-chloro-5-(trifluoromethyl)-2-pyridinyl]oxy]phenyl]methyl]-N'-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

[--00000110]

2/9 - (C) FILE CAPLUS

STN CA Caesar accession number : 1553

AN - 1996:137693 CAPLUS

DN - 124:165248

TI - Aryl antiinflammatory compounds, their preparation, and their activity

IN - Adams, Jerry Leroy; Hall, Ralph Floyd

PA - SmithKline Beecham Corp., USA

SO - PCT Int. Appl., 46 pp.

CODEN: PIXXD2

DT - Patent

LA - English

FAN.CNT 1

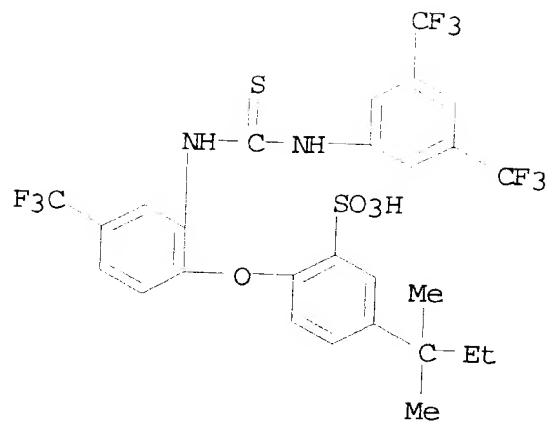
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PN	WO9533458	A	19951214	WO 1995-US6961	19950602
	W: JP, US				
	RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
PR	US 1994-252718		19940602		
OS	MARPAT 124:165248				
AB	The invention relates to the novel compds. and pharmaceutical compns. of I [R1 = SO3H, S(O)n-C1-4 alkyl; n = 0-2; R2 = H, halo, (substituted) C1-8 alkyl, C1-8 alkoxy; m = 1, 2; R3 = C(O)R7, C(S)R7; R4, R8, R9 = H, C1-4 alkyl; R5 = H, halo, CF3, Me, (CH2)tC(O)2R8, (CH2)tOH; t = 0-2; R6 = H, halo; R7 = (substituted) aryl, (substituted) aryl-C1-2 alkyl, (substituted) C1-8 alkyl, NR9R10; R10 = (substituted) aryl, (substituted) aryl-C1-2 alkyl, (substituted) C1-8 alkyl, or R9NR10 form 5- to 7-membered (un)satd. ring with optional addnl. heteroatom of O/N or S; X = O, S; with provisions] and pharmaceutically acceptable salts thereof. The invention also relates to a method of treating or reducing inflammation in a mammal in need thereof, which comprises				

CI 1. Prepn. of selected compds. of the invention is described.  
Compds. of the invention demonstrated phospholipase A2 inhibition,  
generally at 50 .mu.M levels.

AN - 1996:137693 CAPLU  
DN - 124:165248  
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activity  
IN - Adams, Jerry Leroy; Hall, Ralph Floyd  
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FAN.CNT 1

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PR	US 1994-252718		19940602		
OS	MARPAT 124:165248				
IT	---174083-14-8---				
	RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)				
	(aryl antiinflammatory compd. prepn. and activity)				
RN	174083-14-8 CAPLUS				
CN	Benzenesulfonic acid, 2-[2-[[[3,5-bis(trifluoromethyl)phenyl]amino]thioxomethyl]amino]-4-(trifluoromethyl)phenoxy]-5-(1,1-dimethylpropyl)- (9CI) (CA INDEX NAME)				

[--00000018]



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